

ABSTRACT

5 The present invention is directed to novel
replication-deficient adenoviral vectors characterized in
that they harbor at least two lethal early region gene
deletions (E1 and E4) that normally transcribe adenoviral
early proteins. These novel recombinant vectors find
particular use in human gene therapy treatment whereby
10 the vectors additionally carry a transgene or therapeutic
gene that replaces the E1 or E4 regions. The present
invention is further directed to novel packaging cell
lines that are transformed at a minimum with the
adenoviral E1 and E4 gene regions and function to
15 propagate the above novel replication-deficient
adenoviral vectors.

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